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- 1. A purified antibody that preferentially binds a T cell antigen receptor (TCR), wherein said antibody preferentially binds a CDR3-loop or an α - β junction of said TCR; or preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α O⁺ T cells.
- 2. The purified antibody of claim 1, that preferentially binds and preferentially expands an invariant T cell.
- 3. The purified antibody of claim 1, that preferentially binds the antigen binding site of the TCR of said T cell subpopulation.
- 4. A combination of purified antibodies that preferentially binds a TCR, wherein said antibody combination preferentially binds a CDR3-loop or an α - β junction of said TCR; or preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q⁺ T cells; wherein said antibody combination is selected from the group consisting of:
 - (i) an anti-V α 24 antibody and an anti-CD161 antibody;
 - (ii) an anti-Vα24 antibody and an anti-CD94 antibody;
 - (iii) an anti-Vβ11 antibody and an anti-CD161 antibody; and
 - (iv) an anti-Vβ11 antibody and an anti-CD94 antibody.

5. A fragment or derivative of an antibody, wherein said antibody preferentially binds a CDR3-loop or an α - β junction of a TCR; or preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q + T cells.

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- 6. A bifunctional antibody comprising:
- (a) a first antibody or fragment thereof that preferentially binds a CDR3-loop or an α - β junction of a TCR; or preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q ⁺ T cells; wherein said first antibody or fragment binds a first epitope; and
- (b) a second antibody or fragment thereof that binds a second epitope expressed on a T cell expressing said TCR or expressed on a NK T cell, CD1d-reactive T cell, or $J\alpha Q^+$ T cell that is bound by said first antibody or fragment thereof.
- 7. A stable hybridoma that produces an antibody, wherein said antibody preferentially binds a CDR3-loop or an α - β junction of a TCR; or preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q $^+$ T cells.
- 8. A purified T cell subpopulation, wherein said T cells are specifically bound by an antibody or a combination of antibodies, wherein said antibody or said antibody combination preferentially binds a CDR3-loop or an α - β junction of

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a TCR; or wherein said antibody preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and $J\alpha Q^+$ T cells.

- 9. A method of generating an antibody that preferentially binds a CDR3-loop or an α - β junction of a TCR; or preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q ⁺ T cells; said method comprising:
 - (a) coupling a cyclic peptide to a carrier;
 - (b) immunizing a mammal with said coupled peptide; and
- (c) isolating an antibody that preferentially binds a CDR3-loop or an α - β junction of a TCR; or preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q⁺ T cells.
- 10. A method of generating an antibody that preferentially binds a CDR3-loop or an α - β junction of a TCR; or preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q⁺T cells; said method comprising:
- (a) immunizing a CD1 or invariant T cell deficient mammal with invariant T cells; and

(b) isolating an antibody that preferentially binds a CDR3-loop or an α - β junction of a TCR; or preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and $J\alpha Q^+$ T cells.

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- 11. The method of claim 9 or 10, wherein said mammal is a CD1d knockout mouse, a mammal tolerized to NK T cells, a mammal tolerized to CD1d-reactive T cells, a mammal tolerized to $J\alpha Q^+$ T cell, a mammal tolerized to the invariant TCR, a mammal in which invariant T cells have been removed, a mammal lacking part of the α chain of said TCR α chain, or a mammal lacking part of the β chain of said TCR.
- 12. A method of measuring the amount of NK TCRs or the amount of NK T cells in a sample, said method comprising contacting said sample with an antibody that preferentially binds a CDR3-loop, an antigen binding site, or an α - β junction of said TCRs.
- 13. A method of measuring the amount of CD1d-reactive TCRs or the amount of CD1d-reactive T cells in a sample, said method comprising contacting said sample with an antibody that preferentially binds a CDR3-loop, an antigen binding site, or an α - β junction of said TCRs.
- 14. A method of measuring the amount of $J\alpha Q^+$ TCRs or the amount of $J\alpha Q^+$ T cells in a sample, said method comprising contacting said sample with an

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antibody or a combination of antibodies that preferentially binds a CDR3-loop, an antigen binding site, or an α - β junction of said TCRs.

- 15. A method of visualizing the NK TCRs or the NK T cells in a sample,
 said method comprising contacting said sample with an antibody that
 preferentially binds a CDR3-loop, an antigen binding site, or an α-β junction of said TCRs.
 - 16. A method of visualizing the CD1d-reactive TCRs or the CD1d-reactive T cells in a sample, said method comprising contacting said sample with an antibody that preferentially binds a CDR3-loop, an antigen binding site, or an α - β junction of said TCRs.
 - 17. A method of visualizing the $J\alpha Q^+$ TCRs or the $J\alpha Q^+$ T cells in a sample, said method comprising contacting said sample with an antibody or a combination of antibodies that preferentially binds a CDR3-loop, an antigen binding site, or an α - β junction of said TCRs.
- 18. A method of diagnosing a subject with a condition or an increased risk for a condition selected from the group consisting of autoimmune disease, viral infection, bacterial infection, parasitic infection, infection by a eukaryotic pathogen, allergy, asthma, inflammatory condition, graft versus host disease, graft rejection, immunodeficiency disease, spontaneous abortion, pregnancy, and cancer; said method comprising the following:

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- (a) contacting a sample from said subject with an antibody or a combination of antibodies that preferentially binds a CDR3-loop or an α - β junction of a TCR; or an antibody that preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells. CD1d-reactive T cells, and J α Q + T cells;
- (b) quantitating the amount of said antibody or said antibody combination bound to said TCR or said T cells; thereby determining the amount of T cells of interest in said sample; and
- (c) comparing the amount of said T cells of interest in said sample to the amount of said T cells of interest found in subjects diagnosed with said condition or subjects not diagnosed with said condition.
- 19. The method of claim 18, further comprising comparing the amount of another T cell type in said sample with the amount of said another T cell type found in subjects diagnosed with said condition or subjects not diagnosed with said condition.
- 20. A method of treating or preventing an autoimmune disease, viral infection, bacterial infection, parasitic infection, infection by a eukaryotic pathogen, allergy, asthma, inflammatory condition, graft versus host disease, graft rejection, immunodeficiency disease, spontaneous abortion, pregnancy, or cancer in a mammal, said method comprising administering to said mammal an antibody or a combination of antibodies that preferentially binds a CDR3-loop or an α - β junction of a TCR; or preferentially binds or modulates the expansion or

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activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and $J\alpha Q^+$ T cells.

- 21. A method of inhibiting T cell pathogenesis in a mammal, said method
 5 comprising administering to said mammal an antibody or a combination of antibodies that preferentially binds a CDR3-loop, an antigen binding site, or an α-β junction of said TCRs; or inhibits the expansion of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and JαQ+T cells; said administering sufficient to inhibit a T cell expressing said TCR,
 10 a NK T cell, a CD1d-reactive T cell, or a JαQ+T cell.
 - 22. The method of claim 21, wherein said antibody is covalently linked to a toxin or a radiolabel.
 - 23. A method of increasing the size of a subpopulation of T cells, said method comprising contacting a sample comprising said T cells with an antibody that preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, $J\alpha Q^+$ T cells, and T cells expressing a CDR3-loop or an α - β junction of a TCR that is preferentially bound by said antibody, wherein said contacting occurs under conditions that result in an increase in the number of said T cells.
 - 24. The method of claim 23, further comprising contacting said sample with an antigen and antigen presenting cells under conditions that allow said

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contacting to increase the number of said T cells; wherein said antigen is not α -galactosylceramide.

- 25. The method of claim 24, wherein said antigen is
 a lipid or glycosyl-phosphatidylinositol antigen from an infectious pathogen, an antigen from a cancerous cell, or a self-lipid.
 - 26. The method of claim 23, further comprising contacting said sample with an antigen and antigen presenting cells under conditions that allow said contacting to increase the number of said T cells; wherein said antigen is α -galactosylceramide.
 - 27. A method of increasing the size of a subpopulation of T cells, said method comprising:
 - (a) contacting a sample comprising said T cells with an antibody or a combination of antibodies that preferentially binds a CDR3-loop or an α - β junction of a TCR; or an antibody that preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q⁺ T cells; said contacting conducted under conditions that allow complex formation between said T cells and said antibody or said combination of antibodies;
 - (b) isolating said complex; and
 - (c) contacting said T cells in said complex or recovered from said complex with an antigen and antigen presenting cells under conditions that allow said

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contacting to increase the number of said T cells; wherein said antigen is not α -galactosylceramide.

- 28. The method of claim 27, wherein said antigen is a lipid or glycosyl-phosphatidylinositol antigen from an infectious pathogen, an antigen from a cancerous cell, or a self-lipid.
 - 29. A method of increasing the size of a subpopulation of T cells, said method comprising:
 - (a) contacting a sample comprising said T cells with an antibody or a combination of antibodies that preferentially binds a CDR3-loop or an α - β junction of a TCR; or an antibody that preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q $^+$ T cells; said contacting conducted under conditions that allow complex formation between said T cells and said antibody or said combination of antibodies;
 - (b) isolating said complex; and
- (c) contacting said T cells in said complex or recovered from said complex with an antigen and antigen presenting cells under conditions that allow said contacting to increase the number of said T cells; wherein said antigen is wherein said antigen is α-galactosylceramide.

- 30. The method of claim 27 or 29, further comprising contacting said sample or said complex with one or more cytokines.
- 31. A method of increasing the size of a subpopulation of T cells in a mammal, said method comprising:
 - (a) obtaining a sample comprising said T cells from said mammal;
 - (b) contacting said T cells with an antibody or a combination of antibodies that preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, $J\alpha Q^+$ T cells, and T cells expressing a CDR3-loop or an α - β junction of a TCR that is preferentially bound by said antibody or said antibody combination; said contacting conducted under conditions that allow said contacting to increase the number of said T cells; and
 - (c) administering said contacted T cells to said mammal.
 - 32. The method of claim 31, further comprising purifying said T cells prior to said contacting step or after said contacting step.
- 33. A method of increasing the size of a subpopulation of T cells in a mammal, said method comprising:
 - (a) obtaining a sample comprising said T cells from said mammal;
 - (b) contacting said T cells with an antibody or a combination of antibodies that preferentially binds a CDR3-loop or an α - β junction of a TCR; or an antibody that preferentially binds or modulates the expansion or activation of at least one T

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cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and $J\alpha Q^+$ T cells; said contacting conducted under conditions that allow complex formation between said T cells and said antibody or said combination of antibodies;

- 5 (c) isolating said complex; and
 - (d) contacting said T cells in said complex or recovered from said complex with an antigen and antigen presenting cells under conditions that allow said contacting to increase the number of said T cells; wherein said antigen is not α -galactosylceramide; and
 - (e) administering said contacted T cells to said mammal.
 - 34. The method of claim 33, wherein said antigen is a lipid or glycosyl-phosphatidylinositol antigen from an infectious pathogen, an antigen from a cancerous cell, or a self-lipid.
 - 35. A method of increasing the size of a subpopulation of T cells in a mammal, said method comprising:
 - (a) obtaining a sample comprising said T cells from said mammal;
 - (b) contacting said T cells with an antibody or a combination of antibodies that preferentially binds a CDR3-loop or an α - β junction of a TCR; or an antibody that preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and $J\alpha Q^+$ T cells; said contacting conducted under conditions that allow complex

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formation between said T cells and said antibody or said combination of antibodies;

- (c) isolating said complex; and
- (d) contacting said T cells in said complex or recovered from said complex with an antigen and antigen presenting cells under conditions that allow said contacting to increase the number of said T cells; wherein said antigen is α -galactosylceramide; and
 - (e) administering said contacted T cells to said mammal.
- 36. The method of claim 33 or 35, further comprising administering one or more cytokines to said mammal.
 - 37. The method of claim 33 or 35, further comprising contacting said sample or said T cells with one or more cytokines, wherein said contacting alters the ratio of Th1/ Th2/ immune deviation response by said contacted T cells
 - 38. The method of claim 33 or 35, wherein said method is used in the treatment or prevention of an autoimmune disease, viral infection, bacterial infection, parasitic infection, infection by a eukaryotic pathogen, allergy, asthma, inflammatory condition, graft versus host disease, graft rejection, immunodeficiency disease, spontaneous abortion, pregnancy, or cancer in said mammal.

- 39. A method of purifying a subpopulation of T cells from a sample, said method comprising contacting said sample with an antibody or a combination of antibodies that preferentially binds a CDR3-loop or an α - β junction of a TCR; or an antibody that preferentially binds or modulates the expansion or activation of at least one T cell subpopulation selected from the group of NK T cells, CD1d-reactive T cells, and J α Q $^+$ T cells.
- 40. The method of claim 39, further comprising contacting said sample with an anti-V α 24, CD4, CD8, CD56, CD161, or V β 11 antibody.
- 41. The method of claim 39, wherein said antibody is covalently linked to a fluorescent label, wherein said complex is isolated based on the fluorescence signal of said complex.
- 42. The method of claim 39, wherein said antibody is covalently linked to a magnetic label, wherein said complex is isolated based on the magnetism of said complex.

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